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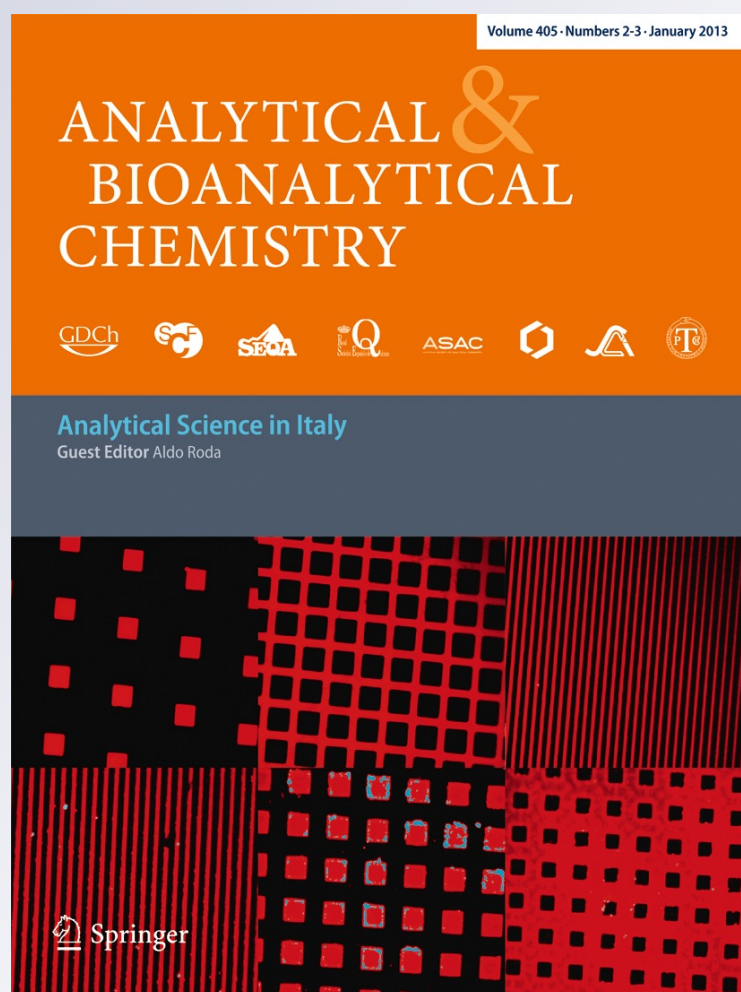
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Speciation analysis of iodine and bromine at picogram-per-gram levels in polar ice

Andrea Spolaor · Paul Vallelonga · Jacopo Gabrieli ·
Natalie Kehrwald · Clara Turetta · Giulio Cozzi ·
Luisa Poto · John M. C. Plane · C. Boutron ·
Carlo Barbante

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Abstract Iodine and bromine species participate in key atmospheric reactions including the formation of cloud condensation nuclei and ozone depletion. We present a novel method coupling a high-performance liquid chromatography with ion chromatography and inductively coupled plasma mass spectrometry, which allows the determination of iodine (I) and bromine (Br) species (IO_3^- , I^- , Br^- , BrO_3^-) at the picogram-per-gram levels presents in Antarctic ice. Chromatographic separation was achieved using an ION-PAC® AS16 Analytical Column with NaOH as eluent. Detection limits for I and Br species were 5 to 9 pg g^{-1} with an uncertainty of less than 2.5% for all considered species. Inorganic iodine and bromine species have been determined in Antarctic ice core samples, with concentrations close to

the detection limits for iodine species, and approximately 150 pg g^{-1} for Br^- . Although iodate (IO_3^-) is the most abundant iodine species in the atmosphere, only the much rarer iodide (I^-) species was present in Antarctic Holocene ice. Bromine was found to be present in Antarctic ice as Br^- .

Keywords Iodine · Bromine · Speciation · Ice cores · Antarctica

Introduction

The atmospheric chemistry of iodine (I) and bromine (Br) are receiving increasing attention due to their importance in

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A. Spolaor
Department of Earth Science, University of Siena,
Via del Laterano 8,
53100 Siena, Italy

A. Spolaor · J. Gabrieli · N. Kehrwald · C. Turetta · G. Cozzi ·
C. Barbante
Institute for the Dynamics of Environmental Processes - CNR,
University of Venice,
Dorsoduro 2137,
30123 Venice, Italy

P. Vallelonga
Centre for Ice and Climate, Niels Bohr Institute,
Juliane Maries Vej 30,
2100 Copenhagen, Denmark

L. Poto · C. Barbante
Department of Environmental Sciences, Informatics and Statistics,
University Ca' Foscari of Venice,
Dorsoduro 2137,
30123 Venice, Italy

J. M. C. Plane
School of Chemistry, University of Leeds,
Leeds LS2 9JT, UK

C. Boutron
Laboratoire de Glaciologie et Géophysique de l'Environnement,
UMR UJF/CNRS 5183,
54, rue Molière, Domaine Universitaire, B.P. 96,
38402 Saint Martin d' Hères, France

C. Barbante (✉)
Accademia Nazionale dei Lincei, Centro B. Segre,
via della Lungara 10,
00196 Rome, Italy
e-mail: barbante@unive.it

insolation-related atmospheric processes [1, 2], formation of cloud condensation nuclei [3], and the chemical depletion of ozone [4]. Furthermore, observations are in discrepancy with model outputs which suggest that iodate should be the main I species in atmospheric depositions [5, 6].

Iodine is a trace element present in the Earth's atmosphere at relatively low concentrations in the range of 1–100 ng m⁻³ [7]. Iodine is principally emitted into the atmosphere in the organic form by oceanic primary production [8], although sea spray aerosol emissions are negligible due to the low concentration of I in seawater (400 to 500 nM [9]). In the ocean, I is mostly present in the IO₃⁻ form [10], even though significant quantities of I⁻ have been detected, most likely as a product of biological cycling [11].

Several chemical forms of I exist in aerosols including the inorganic forms I⁻ and IO₃⁻, and organic iodine compounds (R-I) [12]. Disagreement exists between observations of the atmospheric ratio of I⁻/IO₃⁻, with one study of tropical Atlantic Ocean air finding a majority of IO₃⁻ and negligible I⁻ [13] and another study finding non-negligible I⁻ concentrations in aerosols [14]. Current models of atmospheric chemistry [3] find that the aerosol I⁻ concentration is negligible due to its transformation into species that revert to the gas phase. However, IO₃⁻ is believed to be inert and therefore liable to accumulation with other aerosols [2].

Bromine is a minor constituent in the Earth's atmosphere, but it plays an important role in atmospheric chemistry due to its ozone depletion capabilities [5, 15]. Like iodine, oceans are the main natural reservoir of Br with average concentrations about 65 mg L⁻¹ [16].

Bromine is released into the atmosphere by sea spray aerosol principally as bromide (Br⁻), as well as by formation of volatile bromocarbons from marine algae [17]. It has been observed that the formation of gas phase bromine results in fresh sea salt aerosol being depleted in Br⁻ with respect to sea water [18, 19].

Inorganic and organic speciation of I and Br has been investigated in many environmental matrices including vertical ocean profiles [9], Atlantic coastal waters [20], water macroalgae [21], open ocean deposition [22], and rain [23], as well as snow deposited at the poles [24] and central Europe [5], but no data are available for iodine and bromine species in polar ice. The polar ice caps are important sites for climate studies due to their ability to archive atmospheric constituents, the possibility for accurate sample dating, and their relative distance from anthropogenic pollution sources. Antarctic coastal ice cores additionally offer the possibility to study a wide range of contemporaneous changes in atmospheric and ocean chemistry over several glacial–interglacial cycles [25].

Determination of inorganic I and Br species in polar snow and ice is difficult due to the low concentrations present, and for this reason, the focus of our work has been

to increase the detection sensitivity after species' separation. Determination of I and Br species in water has been achieved with many techniques including ion chromatography (IC) [26] and liquid chromatography (LC) [27] coupled with inductively coupled plasma-mass spectrometry (ICP-MS), UV detection [28], inductively coupled plasma optical emission spectrometry [29], voltammetric techniques [30], and direct analysis of total concentrations by direct ICP-MS analysis [5]. These techniques feature different detection limits with the lowest being reported for IC-LC-ICP-MS (10 to 40 pg g⁻¹) [5] and voltammetric techniques (13 to 25 pg g⁻¹) [30]. Detection limits of UV and polarographic techniques are in the low nanogram-per-gram level [28] which is too high for the determination of I and Br species in polar ice.

Volatile brominated hydrocarbon species have been previously detected [31] in Antarctic ice, but no specific focus was made on the presence of inorganic Br. While inorganic Br in polar ice is likely in the bromide form [5], iodine is mostly present in atmospheric depositions as iodide and iodate; the volatile iodine organic species photo-dissociate rapidly in the atmosphere to generate iodine atoms [32].

Here, we present the first determination of I and Br inorganic species in Antarctic ice core samples. The low concentrations of the investigated species have required the development of a sensitive and reliable method for the speciation analysis of I and Br under ultra-clean conditions at low pictogram-per-gram levels. Using only 0.3 mL of melted ice, samples have been measured by coupling high-performance liquid chromatography (HPLC), with ion chromatography (IC) and inductively coupled plasma mass spectrometry (ICP-MS). ICP-MS is one of the most powerful techniques for rapid analysis of many elements, with low detection limits and high stability compared with other induced plasma techniques such as ICP-AES [33]. The use of ICP-MS, thanks to its multi-elemental capabilities, has also allowed the simultaneous determination of phosphorus species. The novelty of the method allows the use of aggressive eluents with a standard HPLC pump. Interglacial ice samples were analyzed, with particular attention given to the stability of these species after the ice was melted.

Experimental section

Working and standard solutions

All reagent solutions and samples were prepared in a shipping container converted into a particle-free laboratory. The laboratory consists of a series of over-pressured working spaces with air qualities ranging from classes 10 to 100 (US Fed. Std 209E, equivalent to ISO 4 to 5) supplied by modular HEPA fan-filter units (Enviroco, Sanford NC, USA).

Ultrapure water (UPW) was produced by a sequence of active carbon and particulate-filtering cartridges (Millipore, Bedford MA, USA) followed by a mixed bed of ion exchange resins (Maxy, La Gardè, France) [34]. The eluent was prepared by dissolving 1.4 g of NaOH (99.99%, Sigma-Aldrich, Missouri, USA) in 1 L of UPW producing a 35 mM final solution.

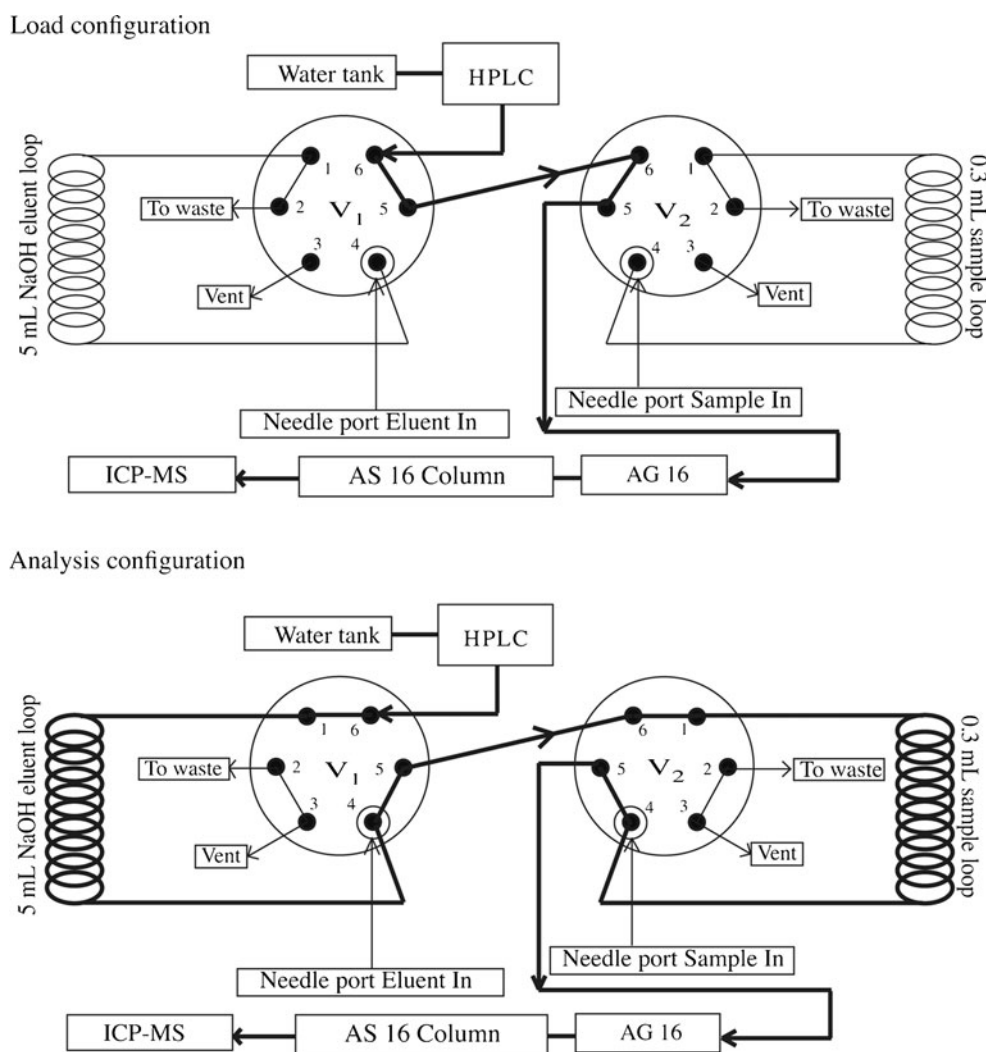
Sample preparation and decontamination

Samples were collected from an ice core drilled at Talos Dome (159°04'21"E, 72°47'14"S 2315 masl) in East Antarctica covered by approximately 1,880 m of ice with an average accumulation rate of $80 \text{ kg m}^{-2} \text{ year}^{-1}$ and a mean annual temperature of -41°C [35]. The samples, dated according to the TALDICE-1 age scale [36, 37], were obtained from a $32 \times 32 \text{ mm}$ square section of the ice core using continuous flow analysis [38]. The ice was melted on a gold-coated brass melt head and distributed to various analytical channels, one of which was used

for discrete sampling. For each 1-m-long ice section melted, an integrated sample of 15 mL was collected. An integrated sample of approximately 15 mL was collected in a Coulter counter accuvette for each metre of ice melted.

To study the stability and potential matrix effects of inorganic iodine and bromine species in melted ice samples, unmelted pieces of Talos Dome ice were analyzed. The unmelted ice core pieces were cut into $1 \times 1.5 \times 3 \text{ cm}$ pieces using a commercial band saw modified with polyethylene tabletop and guides. The table, guides, and saw were carefully cleaned with acetone and methanol before use. A washing procedure [39] was used to remove any surface contamination from the ice core pieces after they had been cut. Samples were decontaminated by immersion in a sequence of three 20°C UPW baths and rinsed with UPW between each immersion step. Samples were immersed in the first bath for 30 to 40 s resulting in the removal of approximately 50% of the sample. The second and third baths of 10 to 15 s removed an additional 20% from the total sample mass. The final sample mass was thus approximately

Fig. 1 A schematic diagram of the sample introduction system in load and analysis configurations. Valve V_1 is controls the eluent loading and V_2 , controls sample introduction. *Thick lines* represent the solution path from the HPLC while *thin lines* represent paths isolated from the HPLC



one third of the initial mass. The decontaminated sample was then kept frozen until analysis to prevent any possible modification in I and Br species.

Apparatus and procedure

The procedure described here for high-resolution determination of I and Br species by HPLC-IC-ICP-MS is based on a system of two valve-controlled loops (Fig. 1), one which handles the NaOH eluent (valve V_1) and the other which handles the sample (valve V_2). Two loops are necessary to allow the use of a stable and reliable HPLC pump while avoiding any damage to the pump by the basic eluent solution (NaOH 35 mM) used. The measurement sequence operates as in the following manner: First the eluent was introduced by V_1 into a 5-mL loop which guaranteed the volume necessary for a single analysis. Second, the melted ice sample was loaded by V_2 into the 0.3-mL loop and injected in the column. Finally, the eluent was flushed through an AG16 guard column and into an AS16 anionic exchange column (Dionex, Milan, Italy). In this way, the columns are able to preconcentrate Br and I species and then separate them with high resolution [5]. The flux into the introduction system was controlled by a HPLC pump, which used UPW as a carrier. The flow was set to 0.280 mL per minute with a run time of 9 min and 70 bar operating pressure. The detector was an Agilent 7500 series ICP-MS (Agilent, CA, USA) with monoisotopic I measured at mass 127 and Br measured at mass 81 (Instrument parameters in Table 1). The ICP-MS data acquisition began automatically when the sample was injected. This analysis is relatively fast and allows the introduction of a large volume of sample (0.3 mL) without the loss of analytes (Table 2) while maintaining low detection limits.

Quantifications

Iodine and bromine species concentrations were quantified by external calibration curves. I and Br standards were prepared from dilutions of 1,000 mg L⁻¹ stock IC standard solutions (TraceCERT® purity grade, Sigma-Aldrich, MO, USA) in UPW. Iodate (IO₃⁻) was prepared by the dissolution of K⁺ salts (TraceCERT® purity grade, Sigma-Aldrich, MO, US) in UPW. No memory effects were present between analyses. Chromatographic data analysis software (Agilent, CA, USA) was used for peak integration.

Results and discussion

Linearity, blanks, and detection limits

Although Antarctic ice can be considered a relatively pure matrix with low elemental concentrations [25], the

Table 1 Instrumental parameters for Agilent 7500 series ICP-MS

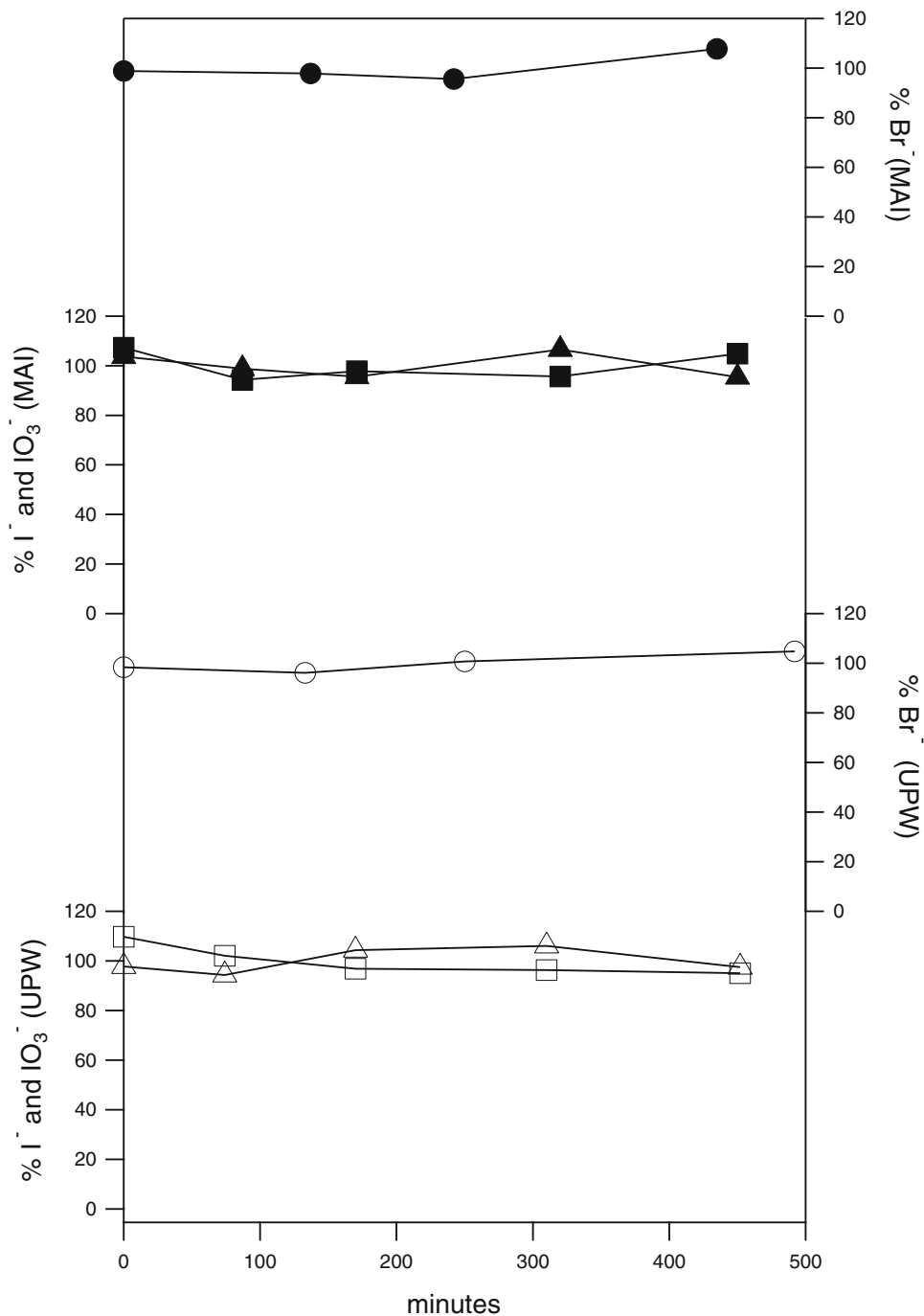
Instrumental parameters	
Plasma parameter	
RF power	1500 W
Sample depth	7.5 mm
Coolant gas	15.0 L/min
Plasma gas	0.75 L/min
Carrier gas	1.20 L/min
Optional gas	0%
Nebulizer pump	0.10 rps
S/C temp	2 degC
Ion lenses	
Extract 1	0.2 V
Extract 2	-121.0 V
Omega-bias-ce	-24 V
Omega-lens-ce	0.4 V
Cell entrance	-32 V
QP focus	-8 V
Cell exit	-44 V
Q-pole parameters	
QP bias	-16.1 V
Octpole parameters	
OctP RF	120 V
OctP bias	-18.0 V
Reaction cell	
H ₂ gas	0 mL/min
He gas	0 mL/min

possibility of matrix effects was nonetheless investigated. An external calibration was conducted as well as a matched calibration, obtained by melting decontaminated ice. Calibrations spanned the ranges 0.07 to 10 ng g⁻¹ for Br⁻ and I⁻ and 0.085 to 13 ng g⁻¹ for IO₃⁻. A matrix effect would be identified as a difference in the gradients of the two calibration curves. For IO₃⁻ and Br⁻, no appreciable differences were observed in the gradients but for I⁻, a matrix effect was present in samples with concentrations greater than 2 ng g⁻¹. For IO₃⁻, the external calibration produced a gradient of 66×10⁴ counts/pg g⁻¹ while the matched calibrations produced a gradient of 64×10⁴ counts/pg g⁻¹. For Br⁻, the external calibration had a value of 11×10⁴ counts/pg g⁻¹ while the matched calibration had a value of 10×10⁴ counts/pg g⁻¹. For I⁻, the external calibration had a gradient of 12×10⁵ counts/pg g⁻¹ while the matched calibration was 10×10⁵ counts/pg g⁻¹. All calibration curves had a correlation coefficient greater than 0.99 (df=3, $p=0.05$). Blank values were obtained by analyzing UPW aliquots according to the same procedure and were used to calculate detection limits. Iodide blank values averaged 3±2 pg g⁻¹, for IO₃⁻ average blank values were 8±2 pg g⁻¹ while for Br⁻ the average blank value was 4±3 pg g⁻¹. The detection limits were

Table 2 Recovery and accuracy of I^- , IO_3^- , and Br^- in an Antarctic ice matrix

	Concentration ($\mu\text{g g}^{-1}$)								
	I^-			IO_3^-			Br^-		
	Added	Measured	Recovery%	Added	Measured	Recovery%	Added	Measured	Recovery%
Sample	0	29 ± 1		0	0	0	0	298 ± 10	0
Addition 1	56	69 ± 2	71	59	65 ± 5	110	50	341 ± 12	86
Addition 2	128	141 ± 4	88	144	130 ± 6	90	123	435 ± 12	111

Fig. 2 Stability of iodine and bromine species in ultrapure water (UPW, *open markers*) and melted Antarctic ice (MAI, *closed markers*) shown as their percentage variations with respect to the mean value. The errors for all species were consistently less than 3%. Circles represent Br^- , squares represent I^- , and triangles represent IO_3^-



calculated as three times the standard deviation of the blanks and ranged from 5 pg g^{-1} for I^- , 7 pg g^{-1} for IO_3^- , and 9 for Br^- . BrO_3^- was not detected in the ice samples, despite the detection limit of our technique being 38 pg g^{-1} . Although IO_3^- was also undetected in the ice samples analyzed here, it has been previously reported in snow deposition [5, 23], and so we have investigated the stability of the species in greater detail.

Recovery, accuracy, and reproducibility

No certified reference material is available for polar ice matrices, therefore we analyzed melted Antarctic samples spiked with different concentrations of I^- , IO_3^- , and Br^- in order to evaluate the accuracy and recovery of the studied species. The concentrations of the three species ranged between 50 and 144 pg g^{-1} (Table 2). Iodide had a weaker retention (71% recovery) at concentrations of 50 pg g^{-1} compared with 88% recovery at 100 pg g^{-1} . Br^- and IO_3^- demonstrated a high recovery for all tested concentrations. To ensure reproducibility, and considering previous studies of iodine speciation [23], concentration of IO_3^- in the matrix-matched standard was fixed to 339 pg g^{-1} . This was required because this species was not found in the interglacial Antarctic ice samples measured. Reproducibility was evaluated by repeating three measurements for each individual sample. All of the determined species demonstrated low residual standard deviations (RSD) of 2.4% for I^- and Br^- and 0.91% for IO_3^- .

Stability of I and Br species in UPW and melted ice

Despite the purity of the Antarctic ice matrix, there remains a possibility that the trace impurities present in that matrix

may alter the species determined here. I and Br species are highly reactive [40], so the stability of these species in samples of melted Antarctic ice was verified under normal laboratory conditions. Figure 2 shows the concentrations of I and Br species determined in a UPW standard over 8 h: No appreciable variations or trends were observed for IO_3^- (open triangles) or Br^- (open circles) over this range of time. I^- (open squares) showed a slight decrease during the period of observation which was most likely due to reaction with O_3 to produce I_2 [13]. To avoid any potential long-term modifications, new I and Br standards were prepared every 2 days. The same experiment was conducted on a melted ice core sample, this time spiked with IO_3^- on account of its absence in Antarctic ice. The melted ice was regularly injected and analyzed every 100 min for 6 h after melting. Our results (closed markers, Fig. 2) are in agreement with the literature [41] and do not show any appreciable change. This study demonstrates that I and Br species are stable in melted ice samples for at least 6 h—much longer than the time required to melt and analyze the Antarctic samples described here.

Ice core samples

A series of interglacial ice core samples were analyzed covering the age between 8 and 12 Ky before present (BP) (Fig. 3). I^- concentrations had a mean value of 11 pg g^{-1} (maximum 31 pg g^{-1}), which was marginally greater than the detection limit of 5 pg g^{-1} . Br^- was found to have a mean concentration of 143 pg g^{-1} and ranged between 92 and 213 pg g^{-1} . Iodate concentrations were always less than the detection limit (7 pg g^{-1}).

These results are consistent with sea salt aerosol being the most important source of Br^- in Antarctic snow [42]. As

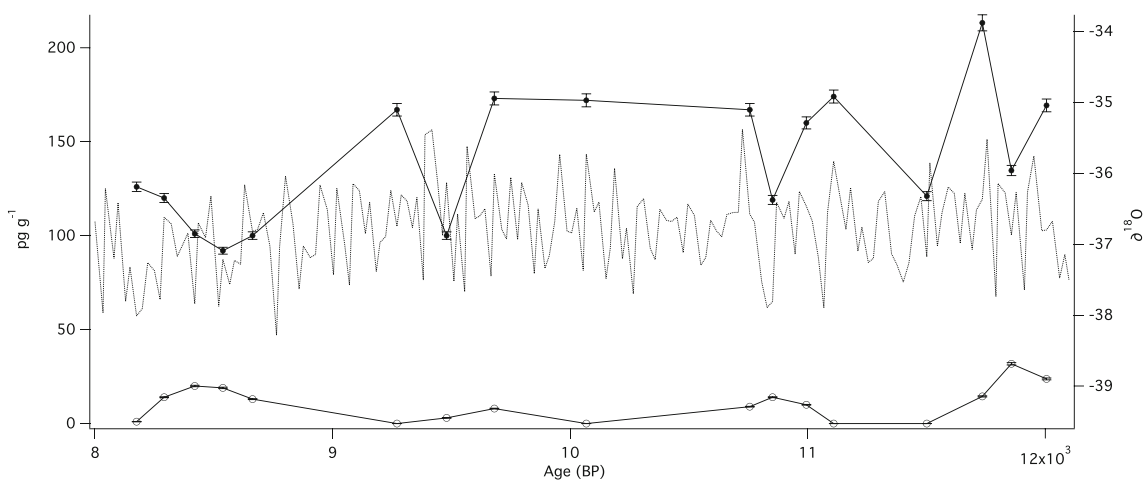


Fig. 3 I^- concentrations (empty circles) and Br^- concentrations (full circles) expressed in pictograms per gram (left axis) in Talos Dome ice core samples, between 8 and 12 thousand years before present (BP).

An isotopic temperature proxy is displayed on the right axis. The measurement uncertainty is smaller than the marker symbol

would be expected for a period of generally stable climate (the paleotemperature proxy $\delta^{18}\text{O}$ [37] is also shown in Fig. 3), Br^- concentrations (Fig. 3, black dots) display little variability.

Iodide concentrations displayed strong variability with occasionally very low values (Fig. 3). Recent studies of marine aerosols [43] and tropical Atlantic Ocean air [22] demonstrated the presence of both iodine oxidation states, IO_3^- and I^- , even though atmospheric chemistry models indicate that atmospheric I should be mainly present in the form of IO_3^- , due to the transformation of I^- into species that revert to the gas phase [44]. Although the ice samples considered in this study originate from a completely different geographic area, our results confirm the findings of Gilfedder et al. [5] who studied I and Br in snowpack. Despite the stability of iodate in particulate and aqueous solutions, only iodide was detected in the ice core samples reported here.

Conclusions

For the first time, iodine and bromine speciation has been investigated in interglacial Antarctic ice core samples. The proposed method is fast and sensitive, with the ability to reach detection limits in the low picogram-per-gram range (5 pg g^{-1} for I^- , 7 pg g^{-1} for IO_3^- , and 9 pg g^{-1} for Br^-) required to quantify the inorganic bromine and iodine species present in Antarctic ice core samples, with an RSD below 3% for all species. The analytical approach adopted here could be useful for many HPLC applications where aggressive eluents are required. Particular attention was paid to contamination control: Sample preparation was carried out in a particle-free laboratory (classes 10 to 100), and the samples were decontaminated by sequential washing.

From our results, the iodine and bromine species analyzed are stable in ultra-pure water and in melted Antarctic ice at least for 6 h—sufficient time for analysis. The interglacial ice core results reported here indicate that both I and Br are present at concentrations in the picogram-per-gram range, but only the inorganic species are present.

Bromide was the only bromine species detected in the analyzed samples and appears not to vary greatly. Iodine was present only in I^- form and displayed greater variability. This may indicate changes in marine primary productivity, since the main source of iodine in the atmosphere is the release of organic compounds from the ocean surface. IO_3^- was not present above detection limits, even though it is believed to be the most stable form of iodine in the atmosphere. Due to the complexity of the atmospheric iodine cycle, further research is necessary to understand the behavior of this species in polar regions. It is also important to investigate the possibility of post-depositional processes which might affect the stability of iodine species in snow and ice.

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